SEQUENCE LISTING

<110> SYNGENTA PARTICIPATIONS AG <120> EXPRESSION IN PLANTS OF ANTIBODIES AGAINST ENTEROTOXIGENIC ESCHERICHIA COLI <130> S 70235/WO <150> US 60/448429 <151> 2003-02-18 <160> 80 <170> PatentIn version 3.1 <210> 1 <211> 399 <212> DNA <213> Artificial Sequence <220> <223> anti-K99 codon optimised VH actagtgagg tgcagctcgt ggagtccggc ggcggcttcg tgaagccggg cggctccctc 60 aageteteet gegeegeete eggetteace tteteegaet actteatgte etggattege 120 cagaeccegg agaagegeet ggagtgggte gecaecatea acaaeggegg eteccaeaee 180 tactgctccg acaacgtgaa gggccgcttc accaccttcc gcgacaacgt gaagaacacc 240 ctctacctcc agatgtcctc cctcaacttc gaggacaccg ccatgtacta ctgcgcccgc 300

gectactacc gettegacgt gegegeetgg ttetectact ggggeeaggg caccetegtg 360

. 399

accgtgtcca cggccaagac cacccgccg tccgtctac

			2			
<210> 2				•		
<211> 58	2					
<212> DN	A					
<213> Ar	tificial Se	quence				
اب د						
<220>	, .					•
<223> and	ti K99 Codo	optimised	VL.			
<400> 2			•			
				•	ccagcgcgtg	
teetteteet	geegegeete	ccagatcato	ggcaccacca	a tccactggto	ccagcagcgc	12
accgacggct	ccccgcgcct	cctcatccag	tgegeetee	g agtccatctc	: cggcatcccg	180
tecegettet	ccggcaccgg	g ctccggcacc	gacttcacco	c tcaacttcaa	ctccgtggag	240
tccgagtaca	ı tcaccgacta	ctactgccag	cagtccaaca	cctggccgac	ctacccgttc	300
ggċggcggca	ccaagetega	gatcaagcgc	gccgacgccg	g ccccgaccgt	gtccatcttc	360
ccgccgțcct	ccgagcaget	cacctccggc	ggcgcgtccg	, tggtgtgctt	cctcaacaac	420
ttctacccga	aggacatcaa	cgtgaagtgg	aagategaeg	gctccgagcg	ccagaacggc	480
gtgctcaact	cctggaccga	ccaggactcc	aaggactcca	cctactccat	gtcctccacc	540
ctcaccctca	ccaaggacga	.gtacgagcgc	cacaactcct	ac	!	582
<210> 3						
<211> 399			•	•		
		•				
<212> DNA			•			
<213> Mou	se		•			
			•			
<400> 3 actagtgaag	tgcaactggt	ggagtctggg	ggaggcttcg	tgaagcctgg	agggtccctg	. 60
				atttcatgtc		120
				ataatggtgg	_	180
				gagacaatgt		240
				ccatgtatta		300
				ggggccaagg		360
	cagccaaaac			555555	J=====================================	399
-						ورر

3	
<210> 4	
<211> 330	
<212> DNA	
<213> Mouse	
<400> 4	
actagtgaca tettgetgae teagteteea gecateetgt etatgattee aagacaaaga	60
gtcagtttct cctgcagggc cagtcagatc attggcacaa ccatacactg gtctcagcaa	120
agaacagatg gttctcctag gcttctcata cagtgtgctt ctgagtctat ctctgggatc	180
cottocaggt ttagtggcac tggatcaggg acagatttta ctcttaactt caacagtgtg	240
gagtctgaat atattacaga ttattactgt caacaaagta atacctggcc aacgtacccg	300
ttcggagggg ggaccaagct cgagataaaa	330
<210> 5 ;	•
<211> 396	
<212> .DNA	
<213> Artificial Sequence	
<220>	
<223> anti-K88 Codon optimised VH from 17_44	
<400> 5	
actagtgacg tgcagctcgt ggagtccggc ggcggcctcg tgcagccggg cggctcccgc	60
aageteteet gegeegeete eggetteace tteteeteet tegeeatgea etgggtgege	120
caggeeccag agaagggeet ggagtgggtg geetacatet eeteeggete cateaceate	180
tactacgccg acaccgtgaa gggccgcttc accgtgtccc gcgacaaccc gaagtccacc	240
ctetteetee agatgaeete eeteegeage gaggaeaeeg ceatgtaeta etgegeeege	300
gacgactacg gctcctccgg ctggtacttc gacgtctggg gcgctggcac cacggtgacc	360
gtgtcctcgg ccaagaccac cccgccgtcc gtctac	396
2210	
<210> 6	
<211> 336	
<212> DNA	

<213> Artificial Sequence

gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc 360 tcctcagcca aaacgacacc cccatctgtc tac 393 <210> 8	<220>	
actagtgaca tegtgatgte ecagteceeg teeteceteg cegtgteege tggegagaag 60 gtcaccatgt cetgeaagte eteccagtee etecteaact ecegeaceeg caagaactac 120 ctegeetggt ateageagaa geegggecag teecegaage teeteateta etgggeetee 240 acceptage ecegeggge eggacegette accegegtgt actactgace ecagtectace 240 etecacatet ecteegtgea ggeggaggac eteggegtgt actactgace ecagtectace 240 acceptate eceteeggea eggtaceaag etegag 336 cettoreae eceteeggea eggtaceaag etegag 336 cettoreae 240 ectoreae ecetteggege eggtaceaag etegag 336 cettoreae ecetteggege eggtaceaag etegag 336 cettoreae ecetteggege eggtaceaag etegag 336 cettoreae ecetteggege eggtaceaag etegag 60 ecetoreae ecetteggege eggtaceaag etegag 60 ecetoreae ecogagee ecetoreae ecetoreae ecogagee ecetoreae ecetoreae ecogagee ecetoreae ecogagee ecetoreae ecogagee eco	<223> anti-K88 Codon optimised VL from 17_44	
gtcaccatgt cctgcaagtc ctcccagtcc ctcctcaact cccgcaccg caagaactac ctcgcctggt atcagcagaa gccgggccag tccccgaagc tcctcatcta ctgggcctcc acccgcgagt ccggcgtgcc ggaccgcttc accggctccg gctccggcac cgacttcacc ctcaccatct cctccgtgca ggcggaggac ctcgccgtgt actactgcac ccagtcctac aacctcctca ccttcggcgc cggtaccaag ctcgag <210> 7 <211> 393 <212> DNA <213> Artificial Sequence <220> <223> anti-K88 Codon optimised VH from 36_41 <400> 7 actagtgagg tccagctgca gcagtctga cctgactag tgaagactgg ggcttcagtg aagatatcct gcaaggcttc tgattactca ctcactgatt actacatgca ctgggtcaag cagagccatg gagagagcct tgagtggatt ggatatatta attttacaa tggtgctact aactacaacc agaagttcaa gggcaaggcc acatttactg tagacacatc ctccagcaca gtctacaatgc agttcaacag cctgacatct gaagactctg cggtctatta ttgtgtaaga gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc tcctcaagcca aaacgacacc cccatctgtc tac <210> 8		
ctegectggt atcageagaa geegggecag teecegaage teeteateta etgggeetee 240 accegegagt eeggegtgee ggacegette aceggeteeg geteeggeae egaetteace 300 accedetet eeteegtgea ggeggaggae eteggegtgt actactgeae ecagteetae 330 accedetetea eetteggege eggtaceaag etegag 336 <210> 7 <211> 393 <212> DNA <211> Artificial Sequence <220> <223> anti-K88 Codon optimised VH from 36_41 <400> 7 actagtgagg teeagetgea geagtetgga eetgaactag tgaagactgg ggetteagtg 60 acagageatg ggagaggeet tgattactea eteactgatt actacatgea etgggteaag 120 cagagecatg gagagageet tgagtggatt ggatatatta atttttacaa tggtgetaet 180 acctacaace agaagtteaa gggeaaggee acatttactg tagacacate etceageaca 240 gtetacatge agtteaacag eetgacate gaagactetg eggtetatta ttgtgtaaga 300 gaageattae taeggaacta tgetatggae taetggggte aaggaacete agteacegte 360 teeteageea aaacgacace eccatetgte tae 393 <210> 8		60
accegegagt ceggegtgee ggacegette aceggeteeg geteeggeae egaetteace 240 cteaceatet ceteegtgea ggeggaggae etegeegtgt actaetgeae ceagteetae 300 aaceteetea cetteggege eggtaceaag etegag 336 <210> 7 <211> 393 <212> DNA <213> Artificial Sequence <220> <223> anti-K88 Codon optimised VH from 36_41 <400> 7 actagtgagg tecagetgea geagtetgga cetgaactag tgaagaetgg ggetteagtg 60 aagatateet geaaggette tgattactea eteactgatt actacatgea etgggteaag 120 cagagecatg gagagacet tgagtggatt ggatatatta attttacaa tggtgetaet 180 aactacaace agaagtteaa gggcaaggee acatttactg tagacacate etecageaca 240 gtetacatge agtteaacag cetgacate gaagaetetg eggtetatta ttgtgtaaga 300 gaagcattae taeggaacta tgetatggae taetggggte aaggaacete agteacegte 360 teetcageca aaacgacace cecatetgte tae 393 <210> 8	gtcaccatgt cetgcaagte eteccagtee etectcaaet eeegcaceeg caagaactae	120
acctected ceteogtgea ggeggaggac eteggeggt actactgeac ceagtectac 300 accetectea cetteggege eggtaceaag etegag 336 <210> 7 <211> 393 <212> DNA <213> Artificial Sequence <220> <223> anti-K88 Codon optimised VH from 36_41 <400> 7 actagtgagg tecagetgea geagtetgga eetgaactag tgaagaetgg ggetteagtg 60 acagageeatg gaagageet tgattactea eteactgat actacatgea etgggteaag 120 cagageeatg gaagageet tgattactea eteactgat attittacaa tggtgetaet 180 acactacaace agaagtteaa gggeaaggee acatttactg tagacacate etecageaca 240 gtetacatge agtteaacag eetgacatet gaagaetetg eggtetatta ttgtgtaaga 300 gaageattae taeggaacta tgetatggae taetggggte aaggaacete agteacegte 360 teeteageea aaacgacace eecatetgte tae 393 <210> 8	ctcgcctggt atcagcagaa gccgggccag tccccgaagc tcctcatcta ctgggcctcc	180
aacctcctca ccttcggcgc cggtaccaag ctcgag 336 <210> 7 <211> 393 <212> DNA <213> Artificial Sequence <220> <223> anti-K88 Codon optimised VH from 36_41 <400> 7 actagtgagg tccagctgca gcagtctgga cctgaactag tgaagactgg ggcttcagtg 60 aagatatcct gcaaggcttc tgattactca ctcactgatt actacatgca ctgggtcaag 120 cagagccatg gagagagcct tgagtggatt ggatatatta atttttacaa tggtgctact 180 aactacaacc agaagttcaa gggcaaggcc acatttactg tagacacatc ctccagcaca 240 gtctacatgc agttcaacag cctgacatct gaagactctg cggtctatta ttgtgtaaga 300 gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc 360 tcctcagcca aaacgacac cccatctgtc tac 393 <210> 8	accegegagt ceggegtgee ggacegette aceggeteeg geteeggeac egaetteace	240
<pre><210> 7 <211> 393 <212> DNA <213> Artificial Sequence <220> <223> anti-K88 Codon optimised VH from 36_41 <400> 7 actagtgagg tocagotgca goagtotgga cotgaactag tgaagactgg ggottcagtg 60 aagatatoot goaaggotto tgattactoa otcactgatt actacatgoa otgggtcaag 120 cagagcoatg gagagagcot tgagtggatt ggatatatta attittacaa tggtgctact 180 aactacaaco agaagttcaa gggcaaggoo acattactg tagacacato otccagcaca 240 gtotacatgo agttcaacag cotgacatot gaagactotg oggtotatta ttgtgtaaga 300 gaagcattac tacggaacta tgotatgga tactggggto aaggaacoto agtcacogto 360 toctcagooa aaacgacaco occatotgto tac 393 <210> 8</pre>	ctcaccatct cctccgtgca ggcggaggac ctcgccgtgt actactgcac ccagtcctac	300
<pre><211> 393 <212> DNA <213> Artificial Sequence <220> <223> anti-K88 Codon optimised VH from 36_41 <400> 7 actagtgagg tccagctgca gcagtctgga cctgaactag tgaagactgg ggcttcagtg 60 aagatatcct gcaaggcttc tgattactca ctcactgatt actacatgca ctgggtcaag 120 cagagccatg gagagagcct tgagtggatt ggatatatta atttttacaa tggtgctact 180 aactacaacc agaagttcaa gggcaaggcc acatttactg tagacacatc ctccagcaca 240 gtctacatgc agttcaacag cctgacatct gaagactctg cggtctatta ttgtgtaaga 300 gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc 360 tcctcagcca aaacgacacc cccatctgtc tac 393 <<100</pre>	aacctcctca ccttcggcgc cggtaccaag ctcgag	336
<pre><212> DNA <213> Artificial Sequence <220> <223> anti-K88 Codon optimised VH from 36_41 <400> 7 actagtgagg tccagctgca gcagtctgga cctgaactag tgaagactgg ggcttcagtg 60 aagatatcct gcaaggcttc tgattactca ctcactgatt actacatgca ctgggtcaag 120 cagagccatg gagagagcct tgagtggatt ggatatatta atttttacaa tggtgctact 180 aactacaacc agaagttcaa gggcaaggcc acatttactg tagacacatc ctccagcaca 240 gtctacatgc agttcaacag cctgacatct gaagactctg cggtctatta ttgtgtaaga 300 gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc 360 tcctcagcca aaacgacacc cccatctgtc tac 393 <<210> 8</pre>	<210> 7	
<pre><213> Artificial Sequence <220> <223> anti-K88 Codon optimised VH from 36_41 <400> 7 actagtgagg tocagetgea geagtetgga cetgaactag tgaagaetgg ggetteagtg 60 aagatateet geaaggette tgattactea etcaetgatt actacatgea etgggteaag 120 cagagecatg gagagageet tgagtggatt ggatatatta attettacaa tggtgetaet 180 aactacaace agaagtteaa gggeaaggee acatttactg tagacacate etceageaca 240 gtetacatge agtteaacag cetgacatet gaagaetetg eggtetatta ttgtgtaaga 300 gaagcattae taeggaacta tgetatggae taetggggte aaggaacete agteaeegte 360 teeteageea aaacgacace eccatetgte tae 393 <210> 8</pre>	<211> 393	
<pre><220> <223> anti-K88 Codon optimised VH from 36_41 <400> 7 actagtgagg tccagctgca gcagtctgga cctgaactag tgaagactgg ggcttcagtg 60 aagatatcct gcaaggcttc tgattactca ctcactgatt actacatgca ctgggtcaag 120 cagagccatg gagagagcct tgagtggatt ggatatatta atttttacaa tggtgctact 180 aactacaacc agaagttcaa gggcaaggcc acatttactg tagacacatc ctccagcaca 240 gtctacatgc agttcaacag cctgacatct gaagactctg cggtctatta ttgtgtaaga 300 gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc 360 tcctcagcca aaacgacacc cccatctgtc tac 393 <<210> 8</pre>	<212> DNA	
<pre><220> <223> anti-K88 Codon optimised VH from 36_41 <400> 7 actagtgagg tccagctgca gcagtctgga cctgaactag tgaagactgg ggcttcagtg 60 aagatatcct gcaaggcttc tgattactca ctcactgatt actacatgca ctgggtcaag 120 cagagccatg gagagagcct tgagtggatt ggatatatta atttttacaa tggtgctact 180 aactacaacc agaagttcaa gggcaaggcc acatttactg tagacacatc ctccagcaca 240 gtctacatgc agttcaacag cctgacatct gaagactctg cggtctatta ttgtgtaaga 300 gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc 360 tcctcagcca aaacgacacc cccatctgtc tac 393 <<210> 8</pre>	<213> Artificial Sequence	
<pre><223> anti-K88 Codon optimised VH from 36_41 <400> 7 actagtgagg tecagetgea geagtetgga cetgaactag tgaagaetgg ggetteagtg 60 aagatateet geaaggette tgattactea etcactgatt actacatgea etgggteaag 120 cagagecatg gagagageet tgagtggatt ggatatatta atttttacaa tggtgetaet 180 aactacaace agaagtteaa gggeaaggee acatttactg tagacacate etceageaca 240 gtetacatge agtteaacag ectgacatet gaagaetetg eggtetatta ttgtgtaaga 300 gaageattae taeggaacta tgetatggae taetggggte aaggaacete agteacegte 360 teeteageca aaacgacac eccatetgte tae 393 <210> 8</pre>	•	
<pre><400> 7 actagtgagg tccagctgca gcagtctgga cctgaactag tgaagactgg ggcttcagtg 60 aagatatcct gcaaggcttc tgattactca ctcactgatt actacatgca ctgggtcaag 120 cagagccatg gagagagcct tgagtggatt ggatatatta atttttacaa tggtgctact 180 aactacaacc agaagttcaa gggcaaggcc acatttactg tagacacatc ctccagcaca 240 gtctacatgc agttcaacag cctgacatct gaagactctg cggtctatta ttgtgtaaga 300 gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc 360 tcctcagcca aaacgacacc cccatctgtc tac 393 <<210> 8</pre>	<220>	
<pre><400> 7 actagtgagg tccagctgca gcagtctgga cctgaactag tgaagactgg ggcttcagtg 60 aagatatcct gcaaggcttc tgattactca ctcactgatt actacatgca ctgggtcaag 120 cagagccatg gagagagcct tgagtggatt ggatatatta atttttacaa tggtgctact 180 aactacaacc agaagttcaa gggcaaggcc acatttactg tagacacatc ctccagcaca 240 gtctacatgc agttcaacag cctgacatct gaagactctg cggtctatta ttgtgtaaga 300 gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc 360 tcctcagcca aaacgacacc cccatctgtc tac 393 <<210> 8</pre>	<223> anti-K88 Codon optimised VH from 36 41	
actagtgagg tecagetgea geagtetgga cetgaactag tgaagaetgg ggetteagtg 60 aagatateet geaaggette tgattaetea eteactgatt actacatgea etggggetaag 120 cagageeatg gagagageet tgagtggatt ggatatatta attittaeaa tggtgetaet 180 aactacaace agaagtteaa gggeaaggee acatttaetg tagacacate eteeageaca 240 gtetacatge agiteaacag eetgacatet gaagaetetg eggtetatta tigtgtaaga 300 gaageattae taeggaacta tgetatggae taetggggte aaggaacete agiteaeegte 360 teeteageea aaacgacace eeeatetgte tae 393 <210> 8	-	
cagagecatg gagagagect tgagtggatt ggatatatta atttttacaa tggtgetaet 180 aactacaace agaagtteaa gggcaaggee acatttactg tagacacate etecageaca 240 gtetacatge agtteaacag eetgacatet gaagaetetg eggtetatta ttgtgtaaga 300 gaageattae taeggaacta tgetatggae taetggggte aaggaacete agteacegte 360 teeteageea aaacgacace eecatetgte tae 393 <210> 8		60
aactacaacc agaagttcaa gggcaaggcc acatttactg tagacacatc ctccagcaca 240 gtctacatgc agttcaacag cctgacatct gaagactctg cggtctatta ttgtgtaaga 300 gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc 360 tcctcagcca aaacgacacc cccatctgtc tac 393 <210> 8	aagatateet geaaggette tgattaetea eteaetgatt aetaeatgea etgggteaag	120
gtctacatgc agttcaacag cctgacatct gaagactctg cggtctatta ttgtgtaaga 300 gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc 360 tcctcagcca aaacgacacc cccatctgtc tac 393 <210> 8	cagagecatg gagagagect tgagtggatt ggatatatta atttttacaa tggtgctact	180
gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc 360 tcctcagcca aaacgacacc cccatctgtc tac 393 <210> 8	aactacaacc agaagttcaa gggcaaggcc acatttactg tagacacatc ctccagcaca	240
tectcageca aaacgacace eccatetgte tae 393	gtctacatgc agttcaacag cctgacatct gaagactctg eggtctatta ttgtgtaaga	300
<210> 8	gaagcattac tacggaacta tgctatggac tactggggtc aaggaacctc agtcaccgtc	360
	tectcageca aaacgacace eccatetgte tac	393
	-210. 9	
	<211> 324	
	<212> DNA <213> Artificial Sequence	

•	
5	-
<pre>_<223> anti-K88 Codon optimised VL from 36_41</pre>	
<400> 8 actagtgaaa atgtgctcac ccagtctcca gcaatcatgt ctgcatctcc aggggaaaag	60
gtcaccatga cctgcagggc cagctcaagt gtaagttccc gttacttgca ctggtaccag	120
cagaagtcag gtgcctcccc caaactctgg atttatagca catccaactt ggcttctgga	180
gtccctgctc gcttcagtgg cagtgggtct gggacctctt actctctcac aatcagcagt	240
griggaggetg aagatgetge cacttattac tgccagcaat acagtggtta cccgtggacg	300
ttcggtggag gcaccaagct cgag	324
<210> 9	
<211> 408	
<212> DNA	
<213> Artificial Sequence	
<220>	•
<223> anti-K88 Codon optimised VH from 7_46	
<400> 9 actagtgaag tgaagcttga ggagtctgga ggaggcttgg tgcaacctgg aggatccatg	60
agactetect gtgttgeete tggatteact tteagtaact actggatgaa etgggteege	60
cagtetecag agaagggget tgagtgggtt getgaaatta, gattgacate taataatttt	120 180
gcaacacatt atgcggagtc tgtgaaaggg aggttcacca tctcaagaga tgattccaaa	240
agtagtgtct acctgcaaat gaacaactta agagctgaag acactggcat ttattactgt	300
accaggeett actaeggtgg taggttette tactggtact tegatgtetg gggegeaggg	360
accaeggtea cegteteete aaccaaaacg acaececeat etgtetae	408
	200
<210> 10	
<211> 324	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> anti-K88 Codon optimised VL from 7_46	

actagtgaaa ttgtgctcac ccagtctcca accaccatgg ctgcatctcc cggggagaag

<400> 10 .

ì

			6			
atcactatc	a cctgcagtg	c cageteaag	t ataagttcc	a attacttgca	ttggtatcag	12
cagaagcca	g gattctccc	c taaactctt	g atttatagg	a catccaatct	ggcttctgga	18
gtcccagtt	c gcttcagtg	g cagtgggtct	gggacctct	t actctctcac	aattggcacc	24
atggaggct	g aagatgttgo	c cacttactac	tgccagcag	g gtaatagtat	accattcacg	300
tteggeteg	g ggacaaagct	: cgag	•			324
<210> 11						
<211> 363	1					
<211> 50.						
<213> Moi		•.				•
(213) MO	.pe					
· <400> 11						
	: tggtggagtc	tgggggaggc	ttagtgcago	ctggagggtc	ccggaaactc	60
tcctgtgcac	cctctggatt	cactttcagt	agetttgcaa	tgcactgggt	tcgtcaggct	120
ccagagaagg	ggctggagtg	ggtcgcatat	attagtagtg	gcagtattac	catctactat	180
gcagacacag	tgaagggccg	attcaccgtc	tccagagaca	atcccaagag	caccctgttc	240
ctgcaaatga	ccagtctaag	gtctgaggac	acggccatgt	attactgtgc	aagagacgac	300
tacggtagta	gcgggtggta	cttcgatgtc	tggggcgcag	ggaccacggt	caccgtctcc	360
tca						363
<210> 12						
<211> 350						
<212> DNA						
<213> Mou	se					
<400> 12	•			•		
gacattgtga	tgtcacagtc	tccatcctcc	ctggctgtgt	cagcaggaga	gaaggtcact	60
atgagetgea	aatccagtca	gagtctgctc	aacagtagaa	cccgaaagaa	ctacttggct	120
tggtaccagc	agaaaccagg	gcagtctcct	aaactgctga	tctactgģgc	atccactagg	180
gaatctgggg	tccctgatcg	cttcacaggc	agtggatctg	ggacagattt	cacyctcacc	240
atcagcagtg	tgcaggctga	agacctggca	gtttattact	gcacgcaatc	ttataatctg	300
ctcacqttcq	gtactagaac	caagetggaa	ctgaatcggg	ctgatgctgc		350

<210> 13 <211> 410 <212> DNA <213> Mouse <400> 13 gaggtccagc tgcagcagtc tggacctgaa ctagtgaaga ctggggcttc agtgaagata 60 tcctgcaagg cttctgatta ctcactcact gattactaca tgcactgggt caagcagagc 120 catggagaga gccttgagtg gattggatat attaattttt acaatggtgc tactaactac 180 aaccagaagt tcaagggcaa ggccacattt actgtagaca catcctccag cacagtctac 240 atgcagttca acagcctgac atctgaagac tctgcggtct attattgtgt aagagaagca 300 ttactacgga actatgctat ggactactgg ggtcaaggaa cctcagtcac cgtctcctca 360 gccaaaacga caccccatc tgtctatcca ctggccccta ctagtgctgc 410 <210> 14 <211> 317 <212> DNA <213> Mouse <400> 14 gaaaatgtgc tcacccagtc tccagcaatc atgtctgcat ctccagggga aaaggtcacc 60 atgacctgca gggccagctc aagtgtaagt tcccgttact tgcactggta ccagcagaag 120 tcaggtgcct cccccaaact ctggatttat agcacatcca acttggcttc tggagtccct 180 gctcgcttca gtggcagtgg gtctgggacc tcttactctc tcacaatcag cagtgtggag 240 gctgaagatg ctgccactta ttactgccag caatacagtg gttacccgtg gacgttcggt 300 ggaggcacca agctgga 317 <210> 15 <211> 374 <212> DNA <213> Mouse

gaagtgaagc ttgaggagtc tggaggaggc ttggtgcaac ctggaggatc catgagactc

<400> 15

	tectgtgttg cetetggatt caettteagt aactaetgga tgaactgggt eegeeagtet	12
	ccagagaagg ggcttgagtg ggttgctgaa attagattga catctaataa ttttgcaaca	18
	cattatgcgg agtctgtgaa agggaggttc accatctcaa gagatgattc caaaagtagt	24
	gtctacctgc aaatgaacaa cttaagagct gaagacactg gcatttatta ctgtaccagg	300
	ccttactacg gtggtaggtt cttctactgg tacttcgatg tctggggcgc agggaccacg	360
	gtcaccgtct cctc	374
	<210> 16	
	<211> 318	•
	<212> DNA	
	<213> Mouse	
	<400> 16	
	gaaattgtgc tcacccagtc tccaaccacc atggctgcat ctcccgggga gaagatcact	60
,	atcacctgca gtgccagctc aagtataagt tccaattact tgcattggta tcagcagaag	120
	ccaggattet cccctaaact ettgatttat aggacateca atetggette tggagtecca	180
	gttegettea gtggeagtgg gtetgggace tettaetete teacaattgg caccatggag	240
	gctgaagatg ttgccactta ctactgccag cagggtaata gtataccatt cacgttcggc	300
	tcggggacaa agctcgag	318
	<210> 17	
	<211> 134	
	<212> PRT ·	
	<213> Artificial Sequence	
	·	
	<220>	
	<223> anti-k99 heavy chain variable region	
	<400> 17	
2	Ala Thr Ser Glu Val Gln Leu Val Glu Ser Gly Gly Phe Val Lys	
-	1 5 10 15	
I	Pro Gly Gly Ser Leu Lys Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe 20 25 30	

Ç

Ser Asp Tyr Phe Met Ser Trp Ile Arg Gln Thr Pro Glu Lys Arg Leu 35 40 45

Glu Trp Val Ala Thr Ile Asn Asn Gly Gly Ser His Thr Tyr Cys Ser 50 55 60

Asp Asn Val Lys Gly Arg Phe Thr Thr Phe Arg Asp Asn Val Lys Asn 65 70 75 80

Thr Leu Tyr Leu Gln Met Ser Ser Leu Asn Phe Glu Asp Thr Ala Met 85 90 95

Tyr Tyr Cys Ala Arg Ala Tyr Tyr Arg Phe Asp Val Arg Ala Trp Phe
100 105 110

Ser Tyr Trp Gly Gln Gly Thr Leu Val Thr Val Ser Thr Ala Lys Thr 115 120 125

Thr Pro Pro Ser Val Tyr

<210> 18

<211> 229

<212> PRT

<213> Artificial Sequence

<220>

<223> anti-k99 light chain

<220>

<221> MISC_FEATURE

<222> (225)..(226)

<223> X at positions 225 and 226 designates an unknown amino acid

<400> 18

Ala Thr Ser Asp Ile Leu Leu Thr Gln Ser Pro Ala Ile Leu Ser Met

1 5 10 15 .

Ile Pro Arg Gln Arg Val Ser Phe Ser Cys Arg Ala Ser Gln Ile Ile
20 25 30

10

Gly Thr Thr Ile His Trp Ser Gln Gln Arg Thr Asp Gly Ser Pro Arg 35 40 45

Leu Leu Ile Gln Cys Ala Ser Glu Ser Ile Ser Gly Ile Pro Ser Arg 50 55 60

Phe Ser Gly Thr Gly Ser Gly Thr Asp Phe Thr Leu Asn Phe Asn Ser 65 70 75 80

Val Glu Ser Glu Tyr Ile Thr Asp Tyr Tyr Cys Gln Gln Ser Asn Thr 85 90 95

Trp Pro Thr Tyr Pro Phe Gly Gly Gly Thr Lys Leu Glu Ile Lys Arg

Ala Asp Ala Ala Pro Thr Val Ser Ile Phe Pro Pro Ser Ser Glu Gln 115 120 125

Leu Thr Ser Gly Gly Ala Ser Val Val Cys Phe Leu Asn Asn Phe Tyr 130 135 140

Pro Lys Asp Ile Asn Val Lys Trp.Lys Ile Asp Gly Ser Glu Arg Gln 145 150 155 160

Asn Gly Val Leu Asn Ser Trp Thr Asp Gln Asp Ser Lys Asp Ser Thr 165 170 175

Tyr Ser Met Ser Ser Thr Leu Thr Leu Thr Lys Asp Glu Tyr Glu Arg

His Asn Ser Tyr Thr Cys Glu Ala Thr His Lys Thr Ser Thr Ser Pro 195 200 205

Ile Val Lys Ser Phe Asn Arg Asn Glu Cys Ser Glu Lys Asp Glu Leu 210 215 220

Xaa Xaa Thr Gly Phe 225

<210> 19

<211> 29

<212> DNA

<213> Artificial Sequence

<220>	11	
<223>	Primer MLALT2	
<400>		
accat	ggatt ttcaagtgca gattttcag	2
<210>	20	
<211>	31	•
<212>	DNA	
<213>	Artificial Sequence	
<220>		
<223>	Primer MLALT3	
<400> caccat	20 ggag wcacakwete agtgtetttr t	31
<210>	21	
<211>	27	
<212>	DNA .	
<213>	Artificial Sequence	
<220>		
<223>	Primer MLALT4	
<400> caccat	21 gkcc ccwrctcagy tyctkgt	27
<210>	22	
<211>	28	
<212>	DNA	
<213>	Artificial Sequence	-
<220>		
	Designer MI ALTE	
4443>	Primer MLALT5	

28

<400> 22

caccatgaag ttgcctgtta ggctgttg

12 <210> 23 <211> 35 <212> DNA <213> Artificial Sequence <220> <223> Primer MH1 <400> 23 atatccacca tggratgsag ctgkgtmats ctctt 35 <210> 24 <211> 35 <212> DNA <213> Artificial Sequence <220> <223> Primer MH2 <400> 24 atatccacca tgracttcgg gytgagctkg gtttt 35 <210> 25 · <211> 30 <212> DNA <213> Artificial Sequence <220> <223> Primer 33615 <400> 25 gaagatctag acttactatg cagcatcagc 30 <210> 26

<211> 27
<212> DNA

<213> Artificial Sequence

13

<220> <223> Primer MVG1R <400> 26 ggcagcacta gtagggcca gtggata 27 <210> 27 <211> 35 <212> DNA <213> Artificial Sequence <220> <223> Primer MVG2R <400> 27 gaggarccac tagtatctcc acacmcaggg gccag 35 <210> 28 <211> 19 <212> PRT <213> Artificial Sequence <220> <223> ER transit peptide <400> 28 Met Arg Val Leu Leu Val Ala Leu Ala Leu Leu Ala Leu Ala Ser Ala Thr Ser <210> 29 <211> 24 <212> DNA

<213> Artificial Sequence

	14	
<220	•	
<223	> Primer	
	> 29 gtcgat catccaggtg caac .	. 24
<210	> 30	
<211:	> 22	
<212>	> DNA	
<213>	Artificial Sequence	
<220>		
<223>	Primer	
<400> actag	30 tggcg ctcgcagcga ga	22
<210>	31	
<211>	25	
<212>	DNA	
<213>	Artificial Sequence	·
<220>		,
<223>	Primer	
<400> accggt	31 tctg ttctgcacaa agtgt	25
<210>	32	
<211>	21 .	
<212>	DNA	
<213>	Artificial Sequence	•
<220>		
<223>	Primer	
<400> acgcgtt	32 Etgt accectggat t	21

15

<210> 33

<211> 22

<212> DNA

<213> Artificial Sequence

<220>

<223> Primer .

<400> 33

acgcgtttgc atgcctgcag tg

<210> 34

<211> 22

<212> DNA

<213> Artificial Sequence

<220>

<223> Primer

<400> 34

agtccaacgg tggagcggaa ct

<210> 35

<211> 6

<212> PRT

<213> Artificial Sequence

<220>

<223> ER retention signal

<400> 35

Ser Glu Lys Asp Glu Leu

<210> 36

<211> 30

<212> DNA

22

<213>	16 Artificial Sequence	
	· •	
<220>		
<223>	Oligonucleotide	
<400> agcttg	36 ggatc cactagtacc ggtacgcgtg	30
<210>	37	
<211>	30	
<212>	DNA	
<213>	Artificial Sequence	
<220>		
<223>	Oligonucleotide	
<400> aattca	37 cgcg taccggtact agtggatcca	30
<210>	38	
<211>	72	
<212>	DNA	
<213>	Artificial Sequence	
	·	
<220>		
<223>	Oligonucleotide	
<400> catgtga	38 agge cacecacaag acetecacet ceccaategt gaagagette aacegeaacg	60
agtgctg	gata ga	72
<210×	39	
<211>	72	
<212>	DNA	
<213>	Artificial Sequence	

<220>

	17	
<223>	Oligonucleotide	
<400> ccggt	39 ctatc agcactegtt geggttgaag etettcaega ttggggaggt ggaggtettg	6
	ggcct ca .	72
<210>	40	
<211>	•	
<212>	·	
	Artificial Sequence	
<220>		
<223>	Oligonucleotide	
<400> agctta	40 .cgcg tggatccact agtgagctcg gtaccg	36
<210>	41	
<211>	36	
<212 ['] >	DNA	
<213>	Artificial Sequence	
<220>		
<223>	Oligonucleotide	
<400> aattcgg	41 gtac cgagctcact agtggatcca cgcgta	36
<210>	42	
<211>	54	
<212>	DNA .	
<213>	Artificial Sequence	
:220>	•	
	Oligonucleotide	
	42	

18 <210> 43 <211> 54 <212> DNA <213> Artificial Sequence <220> <223> Oligonucleotide <400> 43 agettgaatt eggtacettg ageteetate acagetegte etteteggae ttge <210> 44 <211> 23 <212> DNA <213> Artificial Sequence <220> <223> Primer IIA <400> 44 . aagcagtggt atcaacgcag agt 23 <210> 45 <211> 30 <212> DNA <213> Artificial Sequence <220> <223> Primer SMART IIA <400> 45 aagcagtggt atcaacgcag agtacgcggg 30 <210> 46 <211> 22 <212> DNA

<213> Artificial Sequence

<220>		
<223>	Primer K99HC-3'	
<400> aagtag	46 acag atgggggtgt cg	22
<210>	47	٠
<211>	27	
<212>	DNA	
<213>	Artificial Sequence	
<220>		
<223>	Primer K88_746_VAR_H5'	
<400> gccacta	47 agtg aagtgaaget tgaggag _.	27
<210>	48 .	
<211>	26	
<212>	DNA	
<213>	Artificial Sequence	
<220>		
<223>	Primer K88_1744_VAR_H5'	
	48 gtg atgtgcagct ggtgga	26
<210>	49	
<211>	27	
<212>	DNA	
<213>	Artificial Sequence	
<220>		
< 223> 1	Primer K88_3641_VAR_H5'	•
<400>, é	19 gtg aggtccagct gcagcag	27

<212> DNA

	•	
<210>	50 .	
<211>	26	
<212>	DNA	
<213>	Artificial Sequence	
<220>	·	
<223>	Primer K88_746_VAR_L5	
<400>	50 gtga aattgtgctc acccag	20
CCaCca	guga datugugutu deedag	26
<210>	51	
<211>	28	
<212>	DNA	
<213>	Artificial Sequence	
	•	
<220>		
<223>	Primer K88_746_VAR_L3	
<400>	51 . cgag ctttgtcccc gagccgaa	28
		20
<210>	52	
<211>	27	
<212>	DNA	
<213>	Artificial Sequence	
<220>		
<223>	Primer K88_3641_VAR_L5	
<400>	52 gtg aaaatgtget cacccag	27
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	-5-5	۵1
:210>	53	
211>	28	

21 <213> Artificial Sequence <220> <223> Primer K88_3641_VAR_L3 <400> 53 ttatctcgag cttggtgcct ccaccgaa 28 <210> 54 <211> 27 <212> DNA <213> Artificial Sequence <220> <223> Primer K88_1744_VAR_L5 <400> 54 gccactagtg acattgtgat gtcacag 27 <210> 55 · <211> 30 <212> DNA <213> Artificial Sequence <220> <223> Primer K88_1744_VAR_L3 <400> 55 ttatctcgag cttggtccca gcaccgaacg 30 <210> 56 <211> 5 <212> PRT <213> Artificial Sequence <220>

<223> Light Chain Variable Domain Motif

22

<400> 56

Lys Leu Glu Ile Lys 1 5

<210> 57

<211> 972

<212> DNA

<213> Artificial Sequence

<220>

<223> Consensus nucleotide sequence of murine IgG1 Heavy Chain <400> 57 gccaaaacga caccccatc tgtctatcca ctggcccctg gatctgctgc ccaaactaac 60 tecatggtga cectgggatg cetggteaag ggetatttee etgageeagt gaeagtgaee 120 tggaactctg gatccctgtc cagcggtgtg cacaccttcc cagctgtcct gcagtctgac 180 ctctacactc tgagcagetc agtgactgtc ccctccagca cctggcccag cgagaccgtc . 240 acctgeaacg ttgcccaccc ggccagcagc accaaggtgg acaagaaaat tgtgcccagg 300 gattgtggtt gtaagcettg catatgtaca gteecagaag tateatetgt etteatette 360 cccccaaagc ccaaggatgt gctcaccatt actctgactc ctaaggtcac gtgtgttgtg 420 gtagacatca gcaaggatga teeegaggte cagtteaget ggtttgtaga tgatgtggag 480 gtgcacacag ctcagacgca accccgggag gagcagttca acagcacttt ccgctcagtc 540 agtgaacttc ccatcatgca ccaggactgg ctcaatggca aggagttcaa atgcagggta 600 aacagtgcag ctttccctgc ccccatcgag aaaaccatct ccaaaaccaa aggcagaccg 660 aaggetecae aggtgtacae cattecaeet eecaaggage agatggeeaa ggataaagte 720 agtetgaeet geatgataae agaettette eetgaagaea ttaetgtgga gtggeagtgg 780 aatgggcagc cagcggagaa ctacaagaac actcagccca tcatggacac agatggctct 840 tacttcgtct acagcaagct caatgtgcag aagagcaact gggaggcagg aaatactttc 900 acctgetetg tgttacatga gggcetgcac aaccaccata etgagaagag ceteteceae 960 tctcctggta aa 972

<210> 58

<211> 39

<212> DNA

23

<213> Artificial Sequence

<220>

<223> Primer for Human CH3

<400> 58

gaattaagga tecaaageca aaggecagee eegegaace

39

<210> 59

<211> '38

<212> DNA

<213> Artificial Sequence

<220>

<223> Primer for Human CH3

<400> 59

tttattgatt attgctcgag tttacccgga gacaggga

38

<210> 60

<211> 42

<212> DNA

<213> Artificial Sequence

<220>

<223> Primer for Murine CH3

<400> 60

aattaatgaa ttaaggatcc aagaccaagg gccgcccgaa gg

42

<210> 61

<211> 42

<212> DNA

<213> Artificial Sequence

<220>

<223> Primer for Murine CH3

<211> 40

	24	
<400> tttat	61 tgatt attgetegag ettgeeeggg gagtgagaga gg	42
<210>	62	
<211'>	38	
<212>	DNA	•
<213>	Artificial Sequence	
<220>		
<223>	Primer for Bovine CH3	
<400> aattaa	62 atgaa ttaaggatee egeaceaaag geeetgee	38
<210>	63	
<211>	40	•
<212>	DNA	
<213>	Artificial Sequence	
<220>		
<223>	Primer for Bovine CH3	
<400>	63 gatt attgetegag ettgeeggeg gaettggagg	
	gard aregorogay cougetygattygattygatty	40
<210>	64	f
<211>	42	
:212>	DNA	
213>	Artificial Sequence	
:220>		
:223>	Primer for Bovine CH2CH3	•
400> taatga	64 aatt aaggateegg eggeeeatet gtgtteatet te	42
210>	65	

25

<212> DNA

<213> Artificial Sequence

<220>

<223> Primer for Bovine CH2CH3

<400> 65

tttattgatt attgctcgag cttgccggcg gacttggagg

40

<210> 66

<211> 230

<212> PRT

<213> Artificial Sequence

<220>

<223> Consensus amino acid sequence of IgG Fc sequences

<400> 66

His Cys Pro Cys Pro Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Phe 1 5 10 15

Ile Phe Pro Pro Lys Pro Lys Asp Thr Leu Met Ile Ser Arg Thr Pro 20 25 30

Glu Val Thr Cys Val Val Val Asp Val Ser Gln Glu Asp Pro Glu Val 35 40 45

Gln Pro Val Phe Ser Trp Tyr Val Asp Gly Val Glu Val His Thr Ala 50 55 60

Lys Met Leu Thr Lys Pro Arg Glu Glu Gln Phe Asn Ser Thr Tyr Arg 65 70 75 80

Val Val Ser Val Leu Pro Ile Gln His Gln Asp Trp Leu Asn Gly Lys 85 90 95

Glu Phe Lys Cys Lys Val Asn Asn Lys Ala Leu Pro Ala Pro Ile Glu 100 105 110

Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Glu Pro Gln Val

26

Tyr Val Leu Pro Pro Pro Arg Glu Glu Leu Ser Lys Asn Asp Thr Val

Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Pro Asp Ile Ala Val 145 150 155 160

Glu Trp Gln Ser Asn Gly Gln Pro Glu Pro Glu Asn Lys Tyr Lys Thr 165 170 175

Thr Pro Pro Gin Leu Asp Ser Asp Gly Ser Tyr Phe Leu Tyr Ser Lys
180 185 190

Leu Ser Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Thr Phe Thr Cys 195 200 205

Ser Val Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu 210 215 220

Ser Lys Ser Pro Gly Lys 225 230

<210> 67

<211> 471

<212> PRT

<213> Mouse

<400> 67

Thr Met Arg Val Leu Leu Val Ala Leu Ala Leu Leu Ala Leu Ala Ala 1 5 10 15

Ser Ala Thr Ser Asp Val Gln Leu Val Glu Ser Gly Gly Leu Val 20 25 30

Gln Pro Gly Gly Ser Arg Lys Leu Ser Cys Ala Ala Ser Gly Phe Thr 35 40 45

Phe Ser Ser Phe Ala Met His Trp Val Arg Gln Ala Pro Glu Lys Gly 50 55 60

Leu Glu Trp Val Ala Tyr Ile Ser Ser Gly Ser Ile Thr Ile Tyr Tyr 65 70 75 80

Ala Asp Thr Val Lys Gly Arg Phe Thr Val Ser Arg Asp Asn Pro Lys

27

Ser Thr Leu Phe Leu Gln Met Thr Ser Leu Arg Ser Glu Asp Thr Ala .

Met Tyr Tyr Cys Ala Arg Asp Asp Tyr Gly Ser Ser Gly Trp Tyr Phe
115 . 120 125

Asp Val Trp Gly Ala Gly Thr Thr Val Thr Val Ser Ser Ala Lys Thr 130 140

Asn Ser Met Val Thr Leu Gly Cys Leu Val Lys Gly Tyr Phe Pro Glu 165 170 175

Pro Val Thr Val Thr Trp Asn Ser Gly Ser Leu Ser Ser Gly Val His 180 185 190

Thr Phe Pro Ala Val Leu Gln Ser Asp Leu Tyr Thr Leu Ser Ser Ser 195

Val Thr Val Pro Ser Ser Thr Trp Pro Ser Glu Thr Val Thr Cys Asn 210 215 220

Val Ala His Pro Ala Ser Ser Thr Lys Val Asp Lys Lys Ile Val Pro 225 230 235 240

Arg Asp Cys Gly Cys Lys Pro Cys Ile Cys Thr Val Pro Glu Val Ser 245 250 255

Ser Val Phe Ile Phe Pro Pro Lys Pro Lys Asp Val Leu Thr Ile Thr 260 265 . 270

Leu Thr Pro Lys Val Thr Cys Val Val Val Asp Ile Ser Lys Asp Asp 275 280 285

Pro Glu Val Gln Phe Ser Trp Phe Val Asp Asp Val Glu Val His Thr 290 295 300

Ala Gln Thr Gln Pro Arg Glu Glu Gln Phe Asn Ser Thr Phe Arg Ser 305 310 315 320

Val Ser Glu Leu Pro Ile Met His Gln Asp Trp Leu Asn Gly Lys Glu 325 330 335

28

Phe Lys Cys Arg Val Asn Ser Ala Ala Phe Pro Ala Pro Ile Glu Lys 340 345 350

Thr Ile Ser Lys Thr Lys Gly Arg Pro Lys Ala Pro Gln Val Tyr Thr 355 360 365

Ile Pro Pro Pro Lys Glu Gln Met Ala Lys Asp Lys Val Ser Leu Thr 370 . 375 . 380

Cys Met Ile Thr Asp Phe Phe Pro Glu Asp Ile Thr Val Glu Trp Gln 385 390 395 400

Trp Asn Gly Gln Pro Ala Glu Asn Tyr Lys Asn Thr Gln Pro Ile Met 405 410 415

Asp Thr Asp Gly Ser Tyr Phe Val Tyr Ser Lys Leu Asn Val Gln Lys 420 425 430

Ser Asn Trp Glu Ala Gly Asn Thr Phe Thr Cys Ser Val Leu His Glu 435 440 445

Gly Leu His Asn His His Thr Glu Lys Ser Leu Ser His Ser Pro Gly
450 455 460

Lys Ser Glu Lys Asp Glu Leu 465 470

<210> 68

<211> 244

<212> PRT

<213> Mouse

<400> 68

Met Arg Val Leu Leu Val Ala Leu Ala Leu Leu Ala Leu Ala Ser 1 5 10 15

Ala Thr Ser Asp Ile Val Met Ser Gln Ser Pro Ser Ser Leu Ala Val 20 25 30

Ser Ala Gly Glu Lys Val Thr Met Ser Cys Lys Ser Ser Gln Ser Leu 35 40 45

Leu Asn Ser Arg Thr Arg Lys Asn Tyr Leu Ala Trp Tyr Gln Gln Lys 50 55 60

Pro Gly Gln Ser Pro Lys Leu Leu Ile Tyr Trp Ala Ser Thr Arg Glu 65 70 75 80

Ser Gly Val Pro Asp Arg Phe Thr Gly Ser Gly Ser Gly Thr Asp Phe 85 90 95

Thr Leu Thr Ile Ser Ser Val Gln Ala Glu Asp Leu Ala Val Tyr Tyr 100 105 110

Cys Thr Gln Ser Tyr Asn Leu Leu Thr Phe Gly Ala Gly Thr Lys Leu . 115 120 125

Glu Ile Lys Arg Ala Asp Ala Ala Pro Thr Val Ser Ile Phe Pro Pro 130 135 140

Ser Ser Glu Gln Leu Thr Ser Gly Gly Ala Ser Val Val Cys Phe Leu 150 155 160

Asn Asn Phe Tyr Pro Lys Asp Ile Asn Val Lys Trp Lys Ile Asp Gly 165 170 175

Ser Glu Arg Gln Asn Gly Val Leu Asn Ser Trp Thr Asp Gln Asp Ser 180 185 190

Lys Asp Ser Thr Tyr Ser Met Ser Ser Thr Leu Thr Leu Thr Lys Asp 195 200 205

Glu Tyr Glu Arg His Asn Ser Tyr Thr Cys Glu Ala Thr His Lys Thr 210 215 220

Ser Thr Ser Pro Ile Val Lys Ser Phe Asn Arg Asn Glu Cys Ser Glu 225 230 235 240

Lys Asp Glu Leu

<210> 69

<211> 469

<212> PRT

<213> Mouse

<400> 69

30

Met Arg Val Leu Leu Val Ala Leu Ala Leu Leu Ala Leu Ala Ala Ser 1 5 10 15

Ala Thr Ser Glu Val Gln Leu Gln Gln Ser Gly Pro Glu Leu Val Lys 20 25 30

Thr Gly Ala Ser Val Lys Ile Ser Cys Lys Ala Ser Asp Tyr Ser Leu 35 40 45

Thr Asp Tyr Tyr Met His Trp Val Lys Gln Ser His Gly Glu Ser Leu 50 55 60

Glu Trp Ile Gly Tyr Ile Asn Phe Tyr Asn Gly Ala Thr Asn Tyr Asn 65 70 75 80

Gln Lys Phe Lys Gly Lys Ala Thr Phe Thr Val Asp Thr Ser Ser Ser 85 90 95

Thr Val Tyr Met Gln Phe Asn Ser Leu Thr Ser Glu Asp Ser Ala Val

Tyr Tyr Cys Val Arg Glu Ala Leu Leu Arg Asn Tyr Ala Met Asp Tyr 115 ' 120 125

Trp Gly Gln Gly Thr Ser Val Thr Val Ser Ser Ala Lys Thr Thr Pro
130 135 140 ...

Pro Ser Val Tyr Pro Leu Ala Pro Gly Ser Ala Ala Gln Thr Asn Ser 145 150 155 160

Met Val Thr Leu Gly Cys Leu Val Lys Gly Tyr Phe Pro Glu Pro Val 165 170 175

Thr Val Thr Trp Asn Ser Gly Ser Leu Ser Ser Gly Val His Thr Phe 180 185 190

Pro Ala Val Leu Gln Ser Asp Leu Tyr Thr Leu Ser Ser Ser Val Thr 195 200 205

Val Pro Ser Ser Thr Trp Pro Ser Glu Thr Val Thr Cys Asn Val Ala 210 215 . 220

His Pro Ala Ser Ser Thr Lys Val Asp Lys Lys Ile Val Pro Arg Asp 225 230 235 240

Cys Gly Cys Lys Pro Cys Ile Cys Thr Val Pro Glu Val Ser Ser Val 245 250 255

Phe Ile Phe Pro Pro Lys Pro Lys Asp Val Leu Thr Ile Thr Leu Thr 260 265 270

Pro Lys Val Thr Cys Val Val Val Asp Ile Ser Lys Asp Asp Pro Glu 275 280 285

Val Gln Phe Ser Trp Phe Val Asp Asp Val Glu Val His Thr Ala Gln 290 295 300

Thr Gln Pro Arg Glu Glu Gln Phe Asn Ser Thr Phe Arg Ser Val Ser 305 310 315 320

Glu Leu Pro Ile Met His Gln Asp Trp Leu Asn Gly Lys Glu Phe Lys 325 330 335

Cys Arg Val Asn Ser Ala Ala Phe Pro Ala Pro Ile Glu Lys Thr Ile 340 345 350

Ser Lys Thr Lys Gly Arg Pro Lys Ala Pro Gln Val Tyr Thr Ile Pro 355 360 365

Pro Pro Lys Glu Gln Met Ala Lys Asp Lys Val Ser Leu Thr Cys Met 370 375 380

Ile Thr Asp Phe Phe Pro Glu Asp Ile Thr Val Glu Trp Gln Trp Asn 395 400

Gly Gln Pro Ala Glu Asn Tyr Lys Asn Thr Gln Pro Ile Met Asp Thr 405 410 415

Asp Gly Ser Tyr Phe Val Tyr Ser Lys Leu Asn Val Gln Lys Ser Asn 420 425 430

Trp Glu Ala Gly Asn Thr Phe Thr Cys Ser Val Leu His Glu Gly Leu 435 440 445

His Asn His His Thr Glu Lys Ser Leu Ser His Ser Pro Gly Lys Ser 450 455 460

Glu Lys Asp Glu Leu 465

<210> 70

<211> 240

32

<212> PRT

<213> mouse

<400> 70

Met Arg Val Leu Leu Val Ala Leu Ala Leu Leu Ala Leu Ala Ser 1 5 10 15

Ala Thr Ser Glu Asn Val Leu Thr Gln Ser Pro Ala Ile Met Ser Ala 20 25 30

Ser Pro Gly Glu Lys Val Thr Met Thr Cys Arg Ala Ser Ser Ser Val 35 40 45

Ser Ser Arg Tyr Leu His Trp Tyr Gln Gln Lys Ser Gly Ala Ser Pro 50 55 60

Lys Leu Trp Ile Tyr Ser Thr Ser Asn Leu Ala Ser Gly Val Pro Ala 65 70 75 80

Arg Phe Ser Gly Ser Gly Ser Gly Thr Ser Tyr Ser Leu Thr Ile Ser 85 90 95

Ser Val Glu Ala Glu Asp Ala Ala Thr Tyr Tyr Cys Gln Gln Tyr Ser 100 105 110

Gly Tyr Pro Trp Thr Phe Gly Gly Gly Thr Lys Leu Glu Ile Lys Arg

Ala Asp Ala Ala Pro Thr Val Ser Ile Phe Pro Pro Ser Ser Glu Gln 130 135 140

Leu Thr Ser Gly Gly Ala Ser Val Val Cys Phe Leu Asn Asn Phe Tyr 155 150

Pro Lys Asp Ile Asn Val Lys Trp Lys Ile Asp Gly Ser Glu Arg Gln
165 170 175

Asn Gly Val Leu Asn Ser Trp Thr Asp Gln Asp Ser Lys Asp Ser Thr 180 185 190

Tyr Ser Met Ser Ser Thr Leu Thr Leu Thr Lys Asp Glu Tyr Glu Arg

His Asn Ser Tyr Thr Cys Glu Ala Thr His Lys Thr Ser Thr Ser Pro 210 215 220

Ile Val Lys Ser Phe Asn Arg Asn Glu Cys Ser Glu Lys Asp Glu Leu 235 235 240

<210> 71

<211> 474

<212> PRT

<213> Mouse

<400> 71

Met Arg Val Leu Leu Val Ala Leu Ala Leu Ala Leu Ala Leu Ala Ser 1 5 10 15

Ala Thr Ser Glu Val Lys Leu Glu Glu Ser Gly Gly Gly Leu Val Gln
20 25 30

Pro Gly Gly Ser Met Arg Leu Ser Cys Val Ala Ser Gly Phe Thr Phe 35 40 45

Ser Asn Tyr Trp Met Asn Trp Val Arg Gln Ser Pro Glu Lys Gly Leu
50 55 60

Glu Trp Val Ala Glu Ile Arg Leu Thr Ser Asn Asn Phe Ala Thr His 65 70 75 80

Tyr Ala Glu Ser Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asp Ser 85 90 95

Lys Ser Ser Val Tyr Leu Gln Met Asn Asn Leu Arg Ala Glu Asp Thr

Gly Ile Tyr Tyr Cys Thr Arg Pro Tyr Tyr Gly Gly Arg Phe Phe Tyr 115 120 125

Trp Tyr Phe Asp Val Trp Gly Ala Gly Thr Thr Val Thr Val Ser Ser 130 135 140

Ala Lys Thr Thr Pro Pro Ser Val Tyr Pro Leu Ala Pro Gly Ser Ala 145 . 150 . 155 . 160

Ala Gln Thr Asn Ser Met Val Thr Leu Gly Cys Leu Val Lys Gly Tyr 165 170 175 WO 2004/074491 PCT/EP2004/001427 .

34

Phe Pro Glu Pro Val Thr Val Thr Trp Asn Ser Gly Ser Leu Ser Ser 180

Gly Val His Thr Phe Pro Ala Val Leu Gln Ser Asp Leu Tyr Thr Leu 195 200 205

Ser Ser Val Thr Val Pro Ser Ser Thr Trp Pro Ser Glu Thr Val 210 215 220

Thr Cys Asn Val Ala His Pro Ala Ser Ser Thr Lys Val Asp Lys Lys 235 230 235

Ile Val Pro Arg Asp Cys Gly Cys Lys Pro Cys Ile Cys Thr Val Pro 245 250 255

Glu Val Ser Ser Val Phe Ile Phe Pro Pro Lys Pro Lys Asp Val Leu 260 265 270

Thr Ile Thr Leu Thr Pro Lys Val Thr Cys Val Val Val Asp Ile Ser 275 280 285

Lys Asp Asp Pro Glu Val Gln Phe Ser Trp Phe Val Asp Asp Val Glu 290 295 300

Val His Thr Ala Gln Thr Gln Pro Arg Glu Glu Gln Phe Asn Ser Thr 305 310 315 320

Phe Arg Ser Val Ser Glu Leu Pro Ile Met His Gln Asp Trp Leu Asn 325 330 335

Gly Lys Glu Phe Lys Cys Arg Val Asn Ser Ala Ala Phe Pro Ala Pro 340 345 350

Ile Glu Lys Thr Ile Ser Lys Thr Lys Gly Arg Pro Lys Ala Pro Gln 355 360 365

Val Tyr Thr Ile Pro Pro Lys Glu Gln Met Ala Lys Asp Lys Val 370 375 380

Ser Leu Thr Cys Met Ile Thr Asp Phe Phe Pro Glu Asp Ile Thr Val 385 390 395 . 400

Glu Trp Gln Trp Asn Gly Gln Pro Ala Glu Asn Tyr Lys Asn Thr Gln
405 410 415

Pro Ile Met Asp Thr Asp Gly Ser Tyr Phe Val Tyr Ser Lys Leu Asn 420 425 430

Val Gln Lys Ser Asn Trp Glu Ala Gly Asn Thr Phe Thr Cys Ser Val 435 440 445

Leu His Glu Gly Leu His Asn His His Thr Glu Lys Ser Leu Ser His 450 455 460

Ser Pro Gly Lys Ser Glu Lys Asp Glu Leu 465 470

<210> 72

<211> 240

<212> PRT

<213> Mouse

<400> 72

Met Arg Val Leu Leu Val Ala Leu Ala Leu Leu Ala Leu Ala Ser 1 10 15

Ala Thr Ser Glu Ile Val Leu Thr Gln Ser Pro Thr Thr Met Ala Ala 20 25 30

Ser Pro Gly Glu Lys Ile Thr Ile Thr Cys Ser Ala Ser Ser Ile 35 40 45

Ser Ser Asn Tyr Leu His Trp Tyr Gln Gln Lys Pro Gly Phe Ser Pro 50 55 60

Lys Leu Leu Ile Tyr Arg Thr Ser Asn Leu Ala Ser Gly Val Pro Val 65 70 75 80

Arg Phe Ser Gly Ser Gly Ser Gly Thr Ser Tyr Ser Leu Thr Ile Gly 85 90 95

Thr Met Glu Ala Glu Asp Val Ala Thr Tyr Tyr Cys Gln Gln Gly Asn 100 105 110

Ser Ile Pro Phe Thr Phe Gly Ser Gly Thr Lys Leu Glu Ile Lys Arg

Ala Asp Ala Ala Pro Thr Val Ser Ile Phe Pro Pro Ser Ser Glu Gln 130 135 140

36

Leu Thr Ser Gly Gly Ala Ser Val Val Cys Phe Leu Asn Asn Phe Tyr 150 155 160

Pro Lys Asp Ile Asn Val Lys Trp Lys Ile Asp Gly Ser Glu Arg Gln
165 170 175

Asn Gly Val Leu Asn Ser Trp Thr Asp Gln Asp Ser Lys Asp Ser Thr 180 185 190

Tyr Ser Met Ser Ser Thr Leu Thr Leu Thr Lys Asp Glu Tyr Glu Arg

His Asn Ser Tyr Thr Cys Glu Ala Thr His Lys Thr Ser Thr Ser Pro 210 210 225 220

Ile Val Lys Ser Phe Asn Arg Asn Glu Cys Ser Glu Lys Asp Glu Leu 235 235 240

<210> 73

<211> 10

<212> PRT

<213> Artificial Sequence

<220>

<223> Predicted N-terminal sequence of 36/41 VL

<400> 73

Glu Asn Val Leu Thr Gln Ser Pro Ala Ile 1 5 10

<210> 74

<211> 9

<212> PRT

<213> Artificial Sequence

<220>

<223> Obtained N-terminal sequence of 36/41 VL

<400> 74

Val Arg Leu Thr Gln Ser Pro Ala Ile

WO 2004/074491.

```
<210> 75
  <211> 10
  <212> PRT
  <213> Artificial Sequence
 <220>
 <223> Predicted N-terminal sequence of 36/41 VH
 <400> 75
 Glu Val Gln Leu Gln Gln Ser Gly Pro Glu
 <210> 76
 <211> 10
 <212> PRT
 <213> Artificial Sequence
<220>
<223> Obtained N-terminal sequence of 36/41 VH
<400> 76
Glu Val Gln Leu Gln Gln Ser Gly Pro Glu
<210> 77
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Predicted N-terminal sequence of 36/41 derived VL produced in Cor
<400> 77
Glu Asn Val Leu Thr Gln Ser Pro Ala Ile
```

```
<210> 78
  <211> 9
  <212> PRT
  <213> Artificial Sequence
 <220>
 <223> Obtained N-terminal sequence of 36/41 derived VL produced in Corn
 Leu Val Leu Thr Gln Ser Pro Ala Ile
 <210> 79
 <211> 10
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Predicted N-terminal sequence of 36/41 derived VH produced in Cor
<400> 79
Glu Val Gln Leu Gln Gln Ser Gly Pro Glu
<210> 80
<211> 10
<212> PRT
<213> Artificial Sequence
<220>
<223> Obtained N-terminal sequence of 36/41 derived VH produced in Corn
<220>
<221> MISC_FEATURE
<222> (2)..(2)
```

39

<223> X at position 2 denotes that this residue was not obtained during sequencing

<400> 80

Glu Xaa Gln Leu Gln Gln Ser Gly Pro Glu 1 5 10